IN THE CLAIMS

- 1-59. (canceled)
- 60. (currently amended) A hypermutable transgenic mouse wherein <u>germ and somatic</u> cells of said mouse comprise a dominant negative allele of a *PMS2* mismatch repair gene, wherein said dominant negative allele comprises a transation mutation *PMS2-134* allele.
- 61. (currently amended) A hypermutable, transgenic mouse produced by a process comprising the steps of:

introducing a polynucleotide comprising a sequence encoding a dominant negative allele of a *PMS2* mismatch repair gene into a fertilized mouse egg, wherein the dominant negative allele comprises a truncation mutation *PMS2-134* allele, whereby said fertilized mouse egg becomes hypermutable;

implanting the fertilized egg into a pseudopregnant female; and allowing said mouse egg to develop into a hypermutable, transgenic mouse.

- 62. (currently amended) A method of making a hypermutable, fertilized mouse egg comprising introducing into said fertilized mouse egg a polynucleotide comprising a sequence encoding a dominant negative allele of a *PMS2* mismatch repair gene, wherein the dominant negative allele comprises a truncation mutation *PMS2-134* allele, whereby said fertilized mouse egg becomes hypermutable.
 - 63-70. (canceled)
- 71. (currently amended) A method for generating a mutation in a gene of interest comprising the steps of:

introducing a polynucleotide comprising a dominant negative allele of a *PMS2* mismatch repair gene into a fertilized mouse egg, wherein the dominant negative allele comprises a truncation mutation *PMS2-134* allele, whereby the fertilized mouse egg becomes hypermutable;

implanting the fertilized egg into a pseudopregnant female;

allowing said fertilized mouse egg to develop into a hypermutable, transgenic mouse; and testing the mouse to determine whether the gene of interest harbors a mutation.

- 72. (previously presented) The method of claim 71 wherein the step of testing comprises analyzing a nucleotide sequence of the gene of interest.
- 73. (previously presented) The method of claim 71 wherein the step of testing comprises analyzing mRNA transcribed from the gene of interest.
- 74. (previously presented) The method of claim 71 wherein the step of testing comprises analyzing a protein encoded by the gene of interest.
- 75. (previously presented) The method of claim 71 wherein the step of testing comprises analyzing the phenotype of the gene of interest.

76-80. (canceled)

- 81. (previously presented) The method of claim 62 wherein the mismatch repair gene is human PMS2
- 82. (previously presented) The method of claim 81 wherein said mismatch repair gene comprises a truncation mutation at codon 134 as shown in SEQ ID NO:1.
- 83. (previously presented) The method of claim 82 wherein the truncation mutation is a thymidine at nucleotide 424 of wild-type PMS2 as shown in SEQ ID NO:1.
- 84. (previously presented) The hypermutable, transgenic mouse of claim 60 comprising a protein which consists of the first 133 amino acids of human PMS2.

- 85. (previously presented) The hypermutable, transgenic mouse of claim 61 wherein the mismatch repair gene is human PMS2.
- 86. (previously presented) The hypermutable, transgenic mouse of claim 61 wherein the dominant negative allele comprises a truncation mutation at codon 134 as shown in SEQ ID NO:1
- 87. (previously presented) The hypermutable, transgenic mouse of claim 86 wherein the truncation mutation is a thymidine at nucleotide 424 of wild-type PMS2 as shown in SEQ ID NO:1.
 - 88. (new) The mouse of claim 60 wherein the mismatch repair gene is human PMS2.
- 89. (new) The mouse of claim 88 wherein said mismatch repair gene comprises a truncation mutation at codon 134 as shown in SEO ID NO:1.
- (new) The mouse of claim 89 wherein the truncation mutation is a thymidine at nucleotide 424 of wild-type PMS2 as shown in SEO ID NO:1.
 - 91. (new) The method of claim 71 wherein the mismatch repair gene is human PMS2.
- (new) The method of claim 91 wherein said mismatch repair gene comprises a truncation mutation at codon 134 as shown in SEQ ID NO:1.
- (new) The method of claim 92 wherein the truncation mutation is a thymidine at nucleotide 424 of wild-type PMS2 as shown in SEQ ID NO:1.
- 94. (new) The mouse of claim 61 comprising a protein which consists of the first 133 amino acid residues of human PMS2.
- 95. (new) The method of claim 62 wherein the mouse egg comprises a protein which consists of the first 133 amino acid residues of human PMS2.

96. (new) The method of claim 71 wherein said mouse comprises a protein which consists of the first 133 amino acid residues of human PMS2.